

United States Patent [19]**Gagliani et al.**[11] **4,315,077**[45] **Feb. 9, 1982**[54] **POLYIMIDES**[75] Inventors: **John Gagliani; Raymond Lee**, both of San Diego, Calif.[73] Assignee: **International Harvester Company**, Chicago, Ill.[21] Appl. No.: **267,460**[22] Filed: **May 27, 1981****Related U.S. Application Data**

[60] Division of Ser. No. 254,137, Apr. 14, 1981, which is a continuation-in-part of Ser. No. 186,668, Sep. 12, 1980, abandoned.

[51] **Int. Cl.³** **C08J 9/00; C08G 18/16**[52] **U.S. Cl.** **521/77; 521/88; 521/185; 521/189; 528/229; 528/353**[58] **Field of Search** **521/77, 88, 185, 189; 528/229, 353**

[56]

References Cited**U.S. PATENT DOCUMENTS**

Re. 30,213	2/1980	Gagliani et al.	521/185
3,518,219	6/1970	Lavin	521/185
3,726,834	4/1973	Acle, Jr.	521/185
3,966,652	6/1976	Gagliani et al.	521/185
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[57]

ABSTRACT

Terpolyimides prepared from tetracarboxylic acid esters and combinations of heterocyclic, aromatic, and aliphatic diamines and artifacts composed of those terpolymers. Precursors, and methods of converting them to the corresponding terpolymers.

19 Claims, No Drawings

POLYIMIDES

The invention described herein was made in the performance of work under NASA Contract No. NAS9-15484 and is subject to the provisions of Section 305 of the National Aeronautics and Space Act of 1958 (72 Stat. 435; 42 USC 2457).

This application is a division of application Ser. No. 254,137 filed Apr. 14, 1981. Application Ser. No. 254,137 is a continuation-in-part of application Ser. No. 186,668 filed Sept. 12, 1980 (now abandoned).

The present invention relates, in one aspect, to polyimides and, more particularly, to certain novel polyimides which have improved properties by virtue of their being terpolymers derived from tetracarboxylic acids and combinations of heterocyclic, aromatic, and aliphatic diamines.

In other aspects our invention relates to precursors of the just alluded to terpolyimides and their preparation and to the conversion of the precursors to the corresponding terpolymers.

U.S. Pat. No. Re. 30,213 issued Feb. 12, 1980 now abandoned, to John Gagliani et al. for METHOD OF MAKING FOAMED COPOLYIMIDES AND PRODUCTS OBTAINED THEREFROM and U.S. patent application Ser. No. 935,378 filed Aug. 12, 1978, by John Gagliani for POLYIMIDES (now U.S. Pat. No. 4,241,193 issued Dec. 23, 1980) disclose hydrolytically stable copolyimide foams which are fire resistant and give off essentially no smoke or toxic fumes when they are heated to degradation temperatures (application Ser. No. 935,378 was copending with application Ser. No. 186,668). Consequently, those foams are useful in aircraft cabins, space vehicles, and land and sea transport and in a variety of other applications where human life or equipment might be endangered by the overheating of conventional, more flammable, smoke-emitting materials. They can be used, in such applications, in fire containing walls and lightweight structures, to protect fuel tanks and heat sensitive systems, and as void filler materials and thermal, cryogenic, electrical and ocoustical insulations, for example.

We have now discovered that polyimide foams which are in many ways superior to those identified above can be made without sacrificing the desirable attributes of the latter by adding a third diamine of aliphatic character to the precursor from which the polyimide is made. Typical advantages of such foams are increased flexibility and resiliency, greater fatigue resistance, and improved compression set properties.

Compression set is a measure of the extent to which a foam will take on a permanent set or deformation after having been compressed to a stated fraction of its original thickness for a prolonged period of time. This is important in seating applications, for example; materials which are susceptible to compression set reach the point where tactile comfort becomes unacceptable much sooner than those having good compression set properties.

Generally speaking, then, our novel terpolyimides disclosed herein are prepared from precursors which are solutions of a lower alkyl ester of 3,3',4,4'-benzophenonetetracarboxylic acid or a mixture of such esters, an aromatic diamine which is free of aliphatic moieties, a heterocyclic diamine, and an aliphatic diamine. The imide-forming functionalities (the amino and carboxylic

moieties) are preferably present in substantially equimolar amounts.

Exemplary of the aromatic and heterocyclic diamines that can be employed are:

2,6-diaminopyridine
4,4'-diaminodiphenyl sulfone
P,P'-methylene dianiline
4,4'-diaminodiphenyl ether

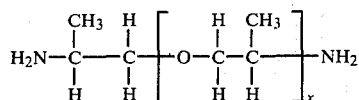
Many other aromatic and heterocyclic diamines have been described in the open and patent literature dealing with the making of polyimides—see, for example, U.S. Pat. Nos. 3,179,614 issued Apr. 20, 1965, to Edwards for POLYAMIDE-ACIDS, COMPOSITIONS THEREOF AND PROCESS FOR THEIR PREPARATION; 3,575,891 issued Apr. 20, 1971, to LeBlanc et al. for STABILIZED POLYIMIDES; and 3,629,180 issued Dec. 21, 1971, to Yoda et al. for PROCESS FOR THE PRODUCTION OF A THERMALLY STABLE POLYMER. Aromatic and heterocyclic diamines selected from those listed in the foregoing patents can be utilized in terpolyimides in accord with the principles of our invention as can others; and we consequently consider our invention to embrace the use of all operable aromatic and heterocyclic diamines.

Aliphatic diamines having from three to 12 carbon atoms have been employed; however, diamines having no more than six carbon atoms will typically prove preferable. The use of those with longer chains can lead to excess thermoplasticity, and that can cause the foam to collapse as it is generated. Also, aliphatic diamines with even numbered chains are preferably employed as they are capable of imparting greater thermal stability to terpolyimides of the character described herein than aliphatic diamines with odd numbered chains.

Aliphatic diamines we have employed include:

1,3-diaminopropane
1,4-diaminobutane
1,6-diaminohexane
1,8-diaminooctane
1,12-diaminododecane
and Jeffamine 230.

The latter is available from the Jefferson Chemical Company and has the formula:



where x is approximately 2.6.

From 0.05 to 0.5 mole of aliphatic diamine per 1.0 mole of benzophenonetetracarboxylic acid can be employed. Concentrations of not more than 0.2 mole per mole of acid are, however, preferred in applications where inflammability is a requirement as the flame resistance of our terpolymers decreases considerably at higher concentrations of the aliphatic diamine.

From 0.05 to 0.9 mole of heterocyclic diamine per mole of acid can be used. Terpolymers with the higher concentration of heterocyclic diamine have the best compression set values and are therefore favored in seat cushioning and other applications of our invention where that property is important.

The precursors of our terpolyimides are essentially monomeric, liquid or solid state solutions of the selected ester (or esters) and diamines.

They are prepared by first reacting 3,3',4,4'-benzophenonetetracarboxylic acid or, preferably, its dianhydride with an esterification agent to form an alkyl diester. Exemplary esterification agents are methyl, ethyl, propyl, and isopropyl alcohols. Methanol is in many cases preferred because of its widespread availability, low cost, and other attributes; because its use facilitates conversion of the precursor to a polyimide foam; and because the foams made from the methyl esters tend to be more flexible, resilient, and compression set resistant. Ethanol is also a preferred esterification agent.

The esterification reaction is followed by the addition of the diamines, which are dissolved in the reaction mixture. The temperature is kept below the reflux temperature of the esterification agent during dissolution of the diamines and low enough to avoid polymerization of the diamines and ester.

Graphite, glass, and other fibers, as well as other fillers such as glass microballoons and additives such as cross-linking agents can be added to the resulting composition to impart wanted properties to the final product. A surfactant can also be added to increase fatigue resistance of the terpolyimide foam and to make it more flexible and resilient by increasing the bubble stability of the foam and the uniformity of the cellular structure.

One preferred surfactant is AS-2, a nonionic, fluorinated, polyalkylene copolymer manufactured by E. I. DuPont de Nemours and Company. We have employed from 0.01 to 0.1 percent of this surfactant based on the weight of the ester and diamine constituents. In systems containing 2,6-diamino pyridine and p,p'-methylene dianiline along with the aliphatic diamine and 3,3',4,4'-benzophenonetetracarboxylic acid ester, a concentration of ca. 0.05 percent proved to be optimum.

Another surfactant that has been successfully employed in those systems in concentrations of 0.1 percent is X-3, a nonionic surfactant of the same general chemical composition as AS-2 and manufactured by the same company.

The material existing after dissolution of the diamines and the addition of any additives may range in form from a "liquid resin" to a spreadable, pastelike formulation depending upon the nature and quantity of any fillers added to the resin. The material may be used in the form just described; or it can be transformed into an amorphous powder capable of being converted into a flexible, resilient, terpolyimide foam. Although not essential, it is preferred that spray drying be employed for this purpose because the liquid resin can thereby be transformed on a continuous basis and in one step into a

dry powder. Also, spray drying allows for modification of the precursor in ways which can be used to vary the properties of the final product.

One suitable spray drying process is described in copending application Ser. No. 186,670 filed Sept. 12, 1980 now U.S. Pat. No. 4,296,208.

The amorphous, powdered resinoid precursor can be converted to a monolithic, terpolyimide foam by various techniques including dielectric, thermal, and microwave heating. The latter, alone or with a thermal postcure, is preferred because of the speed with which the foam can be generated and cured; because the foam is homogeneously heated; and because handling of the fragile, uncured foam can be avoided.

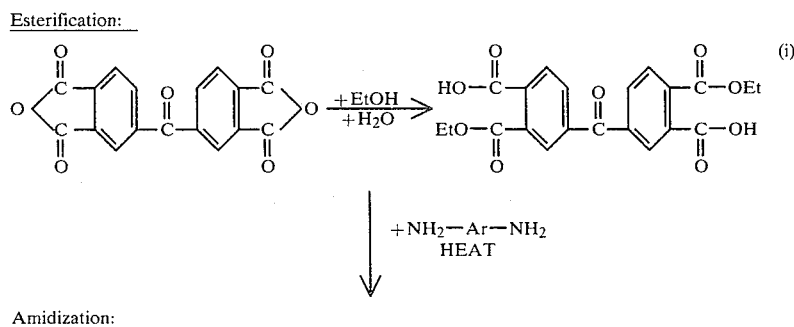
Microwave techniques and equipment that can be used to foam and cure the precursor are disclosed in copending application Ser. No. 186,629 filed Sept. 12, 1980 now U.S. Pat. No. 4,305,796.

Foaming-curing parameters that have proven satisfactory in converting 100 gram samples of representative precursors to flexible, resilient terpolyimide foams are two to 12 minutes exposure to high frequency radiation in an oven operating at a frequency of 2450 MHZ and at 5 kW power followed by thermal heating at a temperature of 500°–550° F. for 15 minutes to two hours.

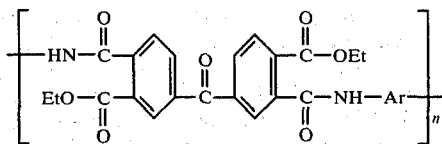
The resulting foam can be employed as such—in a seat cushion or as insulation, for example. Or, using the procedure described in application Ser. No. 935,378 now U.S. Pat. No. 4,241,193 as a further example, the flexible, resilient terpolyimide foam can be converted to a dense, rigid, structurally strong, intumescent material by heating it under pressure.

As suggested above, there are also applications in which the precursor can best be utilized in a liquid or semifluid form. One example is the making of wall and floor panels and other rigid components or artifacts. In a typical application of that character, a layer of the liquid resin, compounded with appropriate fillers, is sandwiched between two pieces of glass cloth wetted with the resin. Foaming and curing of the terpolyimide in a typical wet panel thus formed can be effected in much the same manner as the powdered precursors.

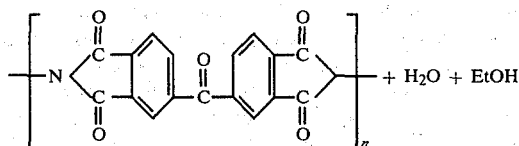
The general model for the chemical reactions which are effected in forming the precursor and in converting it to a polyimide are shown below. The actual reactions are much more complex as three amines, rather than the single aromatic amine shown in the model, are involved



-continued



Imidization:



One advantage of our invention, alluded to above, is that advantage can be taken of its principles to provide polyimides which have the attributes of those state-of-the-art copolyimides disclosed in U.S. Pat. No. Re. 30,213 and in application Ser. No. 935,378, and, at the same time, have such additional desirable attributes as a wider range of mechanical properties, greater flexibility and resilience, and greater fatigue resistance and durability.

Also, our novel compositions have the advantage of great versatility; they can, for example, be produced as foams useful for cushioning and in other applications where comfort is important, and as thermal, electrical, and ocoustical insulations; and they can, on the other hand, be used in floor and wall panels and in other rigid components. They can also be molded into a wide variety of configurations; and fillers and other additives can be compatibly compounded with them to provide optimal performance in various applications of our invention.

Terpolyimides as described above are unique insofar as we are aware. U.S. Pat. Nos. 3,573,132 issued Mar. 30, 1971, to Ducloux et al. for PROCESS FOR THE PREPARATION OF POLYIMIDES and 4,043,978 issued Aug. 23, 1977, to Schmidt et al. for POLYIMIDES suggest that polyimides can be made from tetracarboxylic acids or anhydrides and aliphatic, aromatic, heterocyclic and other diamines and that mixtures of diamines can be employed. However, there is nothing in either patent which would lead one to the particular combination of aromatic plus heterocyclic and aliphatic diamines we employ or which suggests the relative proportions of diamines needed to take advantage of that combination; and there is nothing in either patent which suggests how a polymer with our particular combinations of diamines could be made or that anything would be gained by making such compositions.

From the foregoing it will be apparent to the reader that one important and primary object of our invention resides in the provision of a new family of polymers which are unique in a variety of respects.

Related and also important but more specific objects of the present invention include:

the provision of polymers and compositions containing them which are fire resistant and which give off little or no smoke or toxic compounds when subjected to high temperature, oxidative degradation;

which are strong and durable;

which, when formulated as foams, have improved flexibility and resiliency, fatigue resistance, and compression set;

which are so versatile that they can be employed for such diverse purposes as cushioning and insulation of different types, in panels and other components requiring rigidity and structural strength, and as molding powders.

Another important and again related object of our invention resides in the provision of novel, improved polymers which are terpolyimides derived from a benzophenonetetracarboxylic acid ester and a combination of aromatic, heterocyclic, and aliphatic diamines.

Still other important and primary objects of the present invention reside in the provision of precursors for the polymers identified above and in the provision of processes for making those polymers and for converting the precursors to the corresponding polymers.

Certain important objects of the present invention have been identified above. Other important objects and advantages and additional novel features of the invention will be apparent to those skilled in the relevant arts from the foregoing general description of the invention; from the appended claims; and from the following examples, which are intended to illustrate and not restrict the scope of the invention.

EXAMPLE I

3,3',4,4'-Benzophenonetetracarboxylic acid dianhydride (BTDA) (322.23 g, 1.0 mole) was added to 240 ml of methyl alcohol and 24 ml of water in a one-liter, three-neck flask equipped with a thermometer, a mechanical stirrer, and a reflux condenser. After addition, the mixture was refluxed until clear. The mixture was then refluxed for an additional 60 minutes to ensure complete reaction of the BTDA to its half (or di-) ester.

The contents of the flask were then cooled to 25°-35° C. (77°-95° F.).

2,6-Diaminopyridine (2,6 DAP) (32.8 g, 0.3 mole) and p,p'-methylene dianiline (MDA) (99.1 g, 0.5 mole) were added to the half ester solution and the contents mixed for 15 minutes.

1,6-Diaminohexane (1,6 DAH) (23.7 g, 0.2 mole) was next added to the mixture. This was done slowly enough that the reaction temperature did not exceed 65° C. (149° F.).

The result was a liquid resin precursor which can be used in that form as discussed above.

In a typical instance involving the formation of a low density, high strength, rigid panel, the liquid resin is compounded with selected fillers in a variable speed mixer until the fillers are thoroughly wetted. Glass cloth wetted with the resin is placed on a sheet of aluminum foil. The resin mixture is spread over the glass cloth and covered with another piece of liquid resin wetted glass cloth. Solvent is removed by drying the wet panel in a microwave oven on a sheet of Teflon coated glass cloth at a power output of 1.25 KW for a period of 3 to 5 minutes.

The dried panel is then foamed and cured. Foaming of the panel can be carried out in the microwave oven at a power output of 5.0 KW for six minutes between two sheets of Pyroceran with the thickness of the panel being controlled by Teflon spacers extending between the sheets. The panels can then be cured in a circulating air oven at a temperature of 287.7° C. (550° F.) for 30 minutes.

EXAMPLE II

In an instance leading to a flexible, resilient foam, a liquid resin as described in Example I and made by the process described in that Example was first compounded with 0.1 weight percent of X-3 surfactant, based on the weight of its ester and amino constituents, and then mixed with a 30 phr (parts per hundred parts of resin) dilution ratio of alcohol. A Niro Mobile spray dryer was heated to an inlet temperature of 100° C. (212° F.) and an outlet temperature of 70° C. (158° F.). The liquid resin was then fed into the dryer with the feed being manually adjusted throughout the operation to keep the dryer outlet temperature in the range of 69°-71° C. (156°-160° F.).

This produced a powder which was collected, sieved through a Tyler 48 mesh (297 microns) sieve, and rolled for 30 minutes in a round plastic bottle.

This powder is, essentially, a solid state solution of unreacted diamines and 3,3',4,4'-benzophenonetetracarboxylic acid diester.

A flexible terpolyimide insulating foam was produced from the powder precursor using a Gerling Moore Batch Cavity Model 4115 microwave oven operating at a frequency of 2450 MHz and a power of 5 KW.

The precursor was spread on a Teflon coated glass cloth substrate and placed in the microwave cavity at room temperature. After two to twelve minutes of exposure to the microwave field, depending upon the particular test being conducted, the powder expanded into a homogeneous, cellular foam block. This block was thermally cured into a flexible and resilient foam by heating it at 260° C. (500° F.) for two hours.

The foam rise, cellular structure, resiliency, density, fatigue resistance, and compression set of the foam were than identified.

Resiliency was determined by the ball rebound method described in ASTM Designation D-1564, Suffix B, using a tester fabricated and calibrated in accord with that procedure.

Compression set of the foam at 90 percent compression was determined according to the same ASTM Designation, Method B, using two steel plates held parallel to each other by clamps. The space between the plates was adjusted to the required thickness by spacers.

The resistance of the foam to cycle shear loadings; i.e., its fatigue resistance, was determined in accord with ASTM Designation D-1564, Procedure B, with the exception that examination and measurement of the foam for loss of thickness was made at 10,000 and 20,000 cycles. The fatigue tester was constructed in accord with the same ASTM Designation.

Performance of the foam was detected qualitatively by looking for embrittlement and degradation of the cellular structure and quantitatively by the ball rebound resiliency method and by weight change.

Other tests involved visual observation of the products for cellular structure, foaming behavior, and imperfections; flame resistance using a Meker burner; and hydrolytic stability.

Numerical and qualitative test results are tabulated below:

TABLE 1

Density		90% Compression Set % Loss After 30 Minute Recovery	Resiliency Ball Rebound	Foam Character- istics
lbs/ft ³	kg/m ³			
1.44	23.0	30	55	Flexible, resilient, medium cell size

The foam resisted the open flame of the Meker burner for up to 20 minutes, and it exhibited almost no change after having been kept at 100 percent relative humidity at 60° C. (140° F.) for 30 days.

EXAMPLE III

To demonstrate that other aliphatic diamines can be employed in the novel family of polymers disclosed herein, the procedure described in Example I was repeated, using a variety of aliphatic diamines. The liquid resins thus obtained were then dried and converted to terpolyimide foams using the procedure described in Example II.

Also, to further illustrate how modifications in the formulation of the precursor can be utilized to control the properties of the terpolyimide, the molar ratios of the aliphatic and heterocyclic diamines to the BTDA ester were varied over a considerable range.

The resulting terpolyimide foams were subjected to the analyses described in Example II. Many of the results are tabulated below, and others are discussed in the narrative following the Table:

TABLE 2

Foam Resin Number ³	(Molar Ratios) Aliphatic Diamine	Density		90% Compression Set % Loss After 30 Minute Recovery	Resiliency Ball Rebound	Foam Characteristics
		lbs/ft ³	kg/m ³			
Copolyimides Group 1	None (1.0:0.3:0.6:0.1)	0.538	8.6	52	55	Flexible, resilient, good structure
1	Propyl ²	1.44	23.0	46	50	Flexible, resilient, good structure
2	Butyl	1.32	21.1	63	45	Flexible, resilient, good structure
3	Hexa	1.36	21.8	48	55	Flexible, resilient, good structure
4	Octa	0.943	15.1	39	50	Flexible, resilient, striated

TABLE 2-continued

Foam Resin Number ³	(Molar Ratios) Aliphatic Diamine	Density		90% Compression Set % Loss After 30 Minute Recovery	Resiliency Ball Rebound	Foam Characteristics
		lbs/ft ³	kg/m ³			
5	Dodeca	1.62	25.9	42	50	Flexible, resilient, large cell
6	Jeffamine D-230	1.11	17.8	21	70	size, brittle
Group 2 (1.0:0.2:0.6:0.2)						
7	Propyl	0.840	13.4	40	40	Flexible, resilient, good structure
8	Butyl	1.25	20.0	53	53	Flexible, resilient, good structure
9	Hexa	0.817	13.1	47	55	Flexible, resilient, good structure
10	Octa	1.40	22.4	43	35	Flexible, resilient, good structure
11	Dodeca	3.32	53.0	46	70	Flexible, resilient, good structure
12	Jeffamine D-230	—	—	—	—	Brittle, very large cell size, poor foam
Group 3 (1.0:0.1:0.6:0.3)						
13	Propyl	—	—	—	—	Rigid foam, collapsed and degraded on heating
14	Butyl	1.48	23.7	63	50	Flexible, resilient, fair structure
15	Hexa	1.37	21.9	71	50	Flexible, resilient, fair structure
16	Octa	1.33	21.2	68	45	Flexible, resilient, good structure
17	Dodeca	0.778	13.5	45	70	Flexible, resilient, good structure
Group 4 (1.0:0.3:0.5:0.2)						
18	Propyl	1.33	21.2	40	50	Flexible, resilient, good structure
19	Butyl	0.835	13.4	25	45	Flexible, resilient, good structure
20	Octa	0.845	13.5	22	70	Flexible, resilient, medium cell size
21	Hexa	1.44	23.0	30	55	Flexible, resilient, medium cell size
23	Jeffamine D-230	—	—	—	—	Brittle, very large cell size, collapsed on heating
Group 5 (1.0:0.3:0.4:0.3)						
24	Butyl	1.15	18.3	31	50	Flexible, resilient, good structure
25	Hexa	0.399	6.36	7	55	Flexible, resilient, highly reticulated
Group 6 (1.0:0.3:0.55:0.15)						
26	Hexa	1.17	18.7	44	—	Flexible, resilient, good structure, voids
Group 7 (1.0:0.3:0.65:0.05)						
27	Hexa	1.08	17.3	31	—	Flexible, resilient, good structure, voids

¹In the order of: 3,3',4,4'-benzophenonetetracarboxylic acid ester; 2,6-diamino pyridine; p,p'-methylene dianiline; aliphatic diamine.

²Indication of a radical is used to identify the corresponding aliphatic diamine; e.g., "propyl" = 1,3-diamino propane.

³Each resin contained 0.1 weight percent of X-3 surfactant, and methanol was used as the esterification agent.

⁴This entry, provided for comparison purposes, involved a copolyimide foam derived in essentially the same manner as the foams of Groups 1-7 from a precursor having a 1.0:0.3:0.7 molar ratio of 3,3',4,4'-benzophenonetetracarboxylic acid ester; 2,6-diamino pyridine; and p,p'-methylene dianiline.

The Group 1 foams generally exhibited better compression set values and higher density than foams produced from the copolyimide resin with the Jeffamine D-230 giving useable foams with excellent compression set values (when these foams were scaled up to a large size, the quality of the cell structure worsened).

The Group 2 foams were generally comparable in mechanical characteristics to those of Group 1 except that Jeffamine 230 produced foams which exhibited poor characteristics.

Foams of Group 3 produced smoke and continued to burn for 5-15 seconds after removal from the flame. However, these foams were in many respects satisfactory; and they can accordingly be used where flame resistance is not a controlling criteria.

The Group 4 foams had the most homogeneous cellular structure with the exception of foams made with Jeffamine D-230 (Resin 22). A significant advantage of the foams derived from the precursors of Group 4 is the improved compression set.

the densities of these foams are considerably higher than those of the copolyimide foam which is a draw-

back in applications where weight is at a premium. However, this is offset by increased fatigue resistance.

The Group 5 foams had good structure and excellent compression set properties in one case. However, these foams were found to be less fire resistant than is characteristic of polyimides.

The Group 6 and 7 foams show the effect of varying the concentration of the preferred aliphatic diamine (1,6-diamino hexane). The No. 26 foam was of particular interest. It had considerably decreased fire resistance, indicating that aliphatic diamines concentrations lower than 0.3 mole should be employed in applications where maximum fire resistance is wanted, at least if the aliphatic amine is 1,6 diaminohexane.

EXAMPLE IV

That the properties of our novel terpolyimides can be selectively altered by incorporating a surfactant in the precursor and by varying its concentration is shown by a series of tests conducted as described in Examples I and II except that AS-2 surfactant was employed. The surfactant concentration of the 1.0 BTDA ester; 0.3 2,6 DAP; 0.5 MDA; 0.2 DAH formulation was varied from 0.1 to 1.5 weight percent. The results are tabulated below:

TABLE 3

Foam No.	Surfactant (AS-2)	Density		Indentation Load Deflection (ILD) ¹				Compression Set Loss (Percent)	Resiliency Before Fatigue	After Fatigue (10,000 Cycles)	
	Concentration			N		Lbs.				Resiliency	Height Loss
	Percent	Kg/m ³	Lbs/ft ³	25%	65%	25%	65%			Resiliency	Percent
1	0.1	24.0	1.5	293.6	1427.8	66	321	49	50	55	+3.9

TABLE 3-continued

Surfactant (AS-2)		Indentation Load Deflection (ILD) ¹				Compression	Resiliency	After Fatigue (10,000 Cycles)			
Foam No.	Concentration Percent	Density		N				Lbs.		Set Loss (Percent)	Before Fatigue
		Kg/m ³	Lbs/ft ³	25%	65%	25%	65%				
2	0.25	25.6	1.6	266.9	1352.2	60	304	37	45	47	+5.3
3	0.5	22.4	1.4	266.9	1165.4	60	262	35	40	37	+2.0
4	0.75	22.4	1.4	195.7	1009.7	44	227	34	40	43	-2.9
5	1.0	18.4	1.15	155.7	800.7	35	180	27	45	*	*
6	1.5	17.0	1.06	155.7	809.5	35	182	27	50	*	*

*Cellular structure collapsed after fatigue
¹Performed in accord with ASTM Standard D-1564

EXAMPLE V

We pointed out above that the order in which the diamines are added to the solution of esterified BTDA is an important feature of our invention. This is demonstrated by tests in which the procedures of Examples I and II were followed except for use of the Resin 21 formulation and:

(a) addition of the heterocyclic diamine followed in order by the aromatic and aliphatic diamines at intervals of 15 minutes; and

(b) addition of the aromatic diamine followed in order by the heterocyclic and aliphatic diamines at intervals of 15 minutes.

The addition of the diamines was started with the reaction mixture at a temperature of 30°-35° C. (86°-95° F.). The temperature was allowed to increase freely to approximately 50° C. (122° F.) and then controlled by reducing the rate of the addition of the diamines. Finally, the reaction mixture was heated to and maintained at 60°-65° C. (140°-149° F.) for five minutes.

Addition and complete dissolution of the aromatic diamine before the heterocyclic and the aliphatic diamines were added (option b) produced foams with fewer flaws and significantly less foam collapse at the end of the curing cycle.

EXAMPLE VI

Particle size is another parameter that significantly affects the properties of terpolyimide foams prepared in accord with the principles of our invention. This was demonstrated by a series of tests involving a tepolyimide containing diamino-hexane.

Various particle sizes were obtained by sieving the powdered precursor through a Tyler mesh screen and by comminuting it in a Pulvette bench model grinder.

All foams were produced by foaming and curing the precursor on Teflon coated glass (type 7267/114) in a 15 kW microwave oven using a powder loading of 15 kg (33 lbs.) at a thickness of 6.35 cm (2.5 inches).

The data resulting from this series of tests is summarized below:

TABLE 4

Particle Size (Tyler Mesh)	Indentation Load Deflection				Foam Quality
	25%		65%		
	N	(lbf)	N	(lbf)	
#25	138	39	534	120	Good cellular structure
#50	245	55	1076	242	Rigid structure
Pulverized (maximum size less than 50 microns)	267	60	1054	237	Rigid structure, large flaws

The tabulated data suggests that larger particle size precursors yield more usable foams and a more homogeneous cellular structure with fewer imperfections than foams with a maximum particle size of 50 microns and lower.

The invention may be embodied in other specific forms without departing from the spirit or essential characteristics thereof. The present embodiments are therefore to be considered in all respects as illustrative and not restrictive, the scope of the invention being indicated by the appended claims rather than by the foregoing description; and all changes which come within the meaning and equivalency of the claims are therefore intended to be embraced therein.

What is claimed and desired to be secured by United States Letters Patent is:

1. A method of preparing a terpolyimide foam which includes the steps of: dissolving 3,3',4,4'-benzophenonetetracarboxylic acid or an anhydride thereof in an excess of an alkyl alcohol or mixture of such esters to form a solution of an ester of the acid; adding to and dissolving in said solution in amounts such that the imide forming functionalities are substantially equimolar a heterocyclic diamine, an aromatic diamine, and an aliphatic diamine, there being from 0.05 to 0.5 mole of aliphatic diamine and from 0.05 to 0.9 mole of heterocyclic diamine per mole of 3,3',4,4'-benzophenonetetracarboxylic acid ester; and converting the material existing after said dissolution of said diamines to a polymeric foam.

2. A method of preparing a terpolyimide foam as defined in claim 1 in which said material is converted to a polymeric foam by exposing it to microwave radiation.

3. A method of preparing a terpolyimide foam as defined in claim 2 wherein the exposure of said material to microwave radiation is followed by a thermal post cure.

4. A method of preparing a terpolyimide foam as defined in claim 1 in which said material is converted to a polymeric foam by exposing it to thermal energy.

5. A method of preparing a terpolyimide foam as defined in claim 1 in which said aromatic diamine is added first to said solution and in which said aromatic diamine is fully dissolved before said heterocyclic and aliphatic diamines are added.

6. A method of preparing a terpolyimide foam as defined in claim 1 which includes the step of adding a surfactant to the ester-diamine solution in an amount effective to improve the bubble stability of and produce a more homogeneous cellular structure in the terpolyimide foam to which the precursor is converted.

7. A method of preparing a terpolyimide foam as defined in claim 1 which includes the step of reducing the material existing after the dissolution of the diamines

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in the esterification agent to a dry particulate form before it is converted to a polymeric foam.

8. A method of preparing a terpolyimide foam as defined in claim 1 in which the temperature of the 3,3',4,4'-benzophenonetetracarboxylic acid solution is kept below the boiling point of the esterification agent while the diamines are dissolved in said solution.

9. A method of preparing a terpolyimide foam as defined in claim 1 in which the aromatic diamine is completely dissolved in the material existing after the dissolution of the diamines in the esterification agent before the heterocyclic and aliphatic diamines are added to said material.

10. A method of preparing a terpolyimide foam as defined in claim 1 in which the material existing after the dissolution of the diamines in the esterification agent is reduced to a dry particulate form by spray drying before it is converted to a polymeric foam.

11. A method of preparing a terpolyimide foam as defined in claim 1 which includes the step of intimately admixing a filler with said mixture of diamines and tetracarboxylic acid ester(s).

12. A method of preparing a terpolyimide foam which includes the step of converting to polymeric form with foaming a precursor comprising an essentially stoichiometric mixture of an alkyl ester of 3,3',4,4'-benzophenonetetracarboxylic acid or a mixture of such esters and at least three diamines, one of said diamines being heterocyclic, a second of said diamines being aromatic, a third of said diamines being aliphatic, and said precursor containing from 0.05 to 0.5 mole of aliphatic diamine and from 0.05 to 0.9 mole of heterocyclic diamine per mole of benzophenonetetracarboxylic acid ester(s).

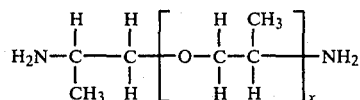
13. A method of preparing a terpolyimide foam as defined in claim 12 in which the aliphatic diamine in the precursor has from three to twelve carbon atoms.

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14. A method of preparing a terpolyimide foam as defined in claim 13 in which the aliphatic diamine in the precursor has not more than six carbon atoms.

15. A method of preparing a terpolyimide foam as defined in either of the preceding claims 13 or 14 in which the aliphatic diamine has an even number of carbon atoms.

16. A method of preparing a terpolyimide foam as defined in claim 12 in which the aliphatic diamine in the precursor has the formula:



where x is on the order of 2.6.

17. A method of preparing a terpolyimide foam as defined in claim 12 in which the precursor is a dry powder and has a maximum particle size of at least 50 microns.

18. A method of preparing a terpolyimide foam as defined in claim 12 in which any ester present in the precursor is a methyl or ethyl half ester of 3,3',4,4'-benzophenonetetracarboxylic acid and the diamines are 2,6-diamino pyridine, p,p'-methylene dianiline, and 1,6-diamino hexane.

19. A method of preparing a terpolyimide foam as defined in claim 18 in which the molar ratio of the ester and diamine constituents in the precursor is:

3,3',4,4'-benzophenonetetracarboxylic acid ester(s)	1.0
2,6-diamino pyridine	0.3
p,p'-methylene dianiline	0.4-0.65
1,6-diamino hexane	0.05-0.3.

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